WEST Search History

Hide Items Restore Clear Cancel

DATE: Monday, March 28, 2005

Hide?	Set Name	<u>e Query</u>	Hit Count
	DB=PG	PB,USPT,EPAB; PLUR=YES; OP=ADJ	
	L13	13 and 19	0
	L12	19 and L1 1	0
	L11	17.ab.	400
	L10	L9 and 17	28
	L9	(536/23.1)![CCLS]	11636
	L8	L7 and l1	1
	L7	aldose reducta\$	1684
	L6	aldose reducta?	1
	L5	TTGTGGCCCACTTCCAGATCGAGAAGCTCT	0
	L4	L3 and 11	1
	L3	L2.ab.	48
	L2	ARL	1090
	L1	dai.in.	2665

END OF SEARCH HISTORY

	FILE	'MEDL]	NE	' ENTE	RED AT	13:4	13:08	ON	28	MAR	2005
L1		0	S	DAI/AU	•						
L2		2321	S	ALDOSE	REDUC	TASE					
L3		312	S	ARL							
L4		2625	S	L2 OR	L3						
L5		0	S '	TTGTGG	CCCACT	TCCAG	SATCG	AGA/	AGC!	ГСТ	
				•							
	FILE	' PCTFU	JLL	' ENTE	RED AT	13:4	13:59	ON	28	MAR	2005
L6	FILE				RED AT						2005
L6 L7	FILE	0	s '		CCCACT						2005
	FILE	0 420	S S	TTGTGG DAI/AU	CCCACT	TCCAG					2005
L7	FILE	0 420 602	s : s :	TTGTGG DAI/AU	CCCACT REDUC	TCCAG					2005
L7 L8	FILE	0 420 602	S : S : S :	TTGTGG DAI/AU ALDOSE L8 AND	CCCACT REDUC	TCCAG					2005

.





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> Search History will be lost after eight hours of inactivity Search numbers may not be continuous; all searches are represented Click on query # to add to strategy To combine searches use # before search number, e.g., #2 AND #6 for |#5 AND #1 Nucleotide History Protein Clipboard Genome <

Details

#6 Search #5 AND #1 Field: Title/Abstract **Most Recent Queries**

#5 Search #4 OR #3 Field: Title/Abstract #4 Search ARL Field: Title/Abstract

#3 Search aldose reductase Field: Title/Abstract

#1 Search dai Field: Author #2 Search aldose reductase Field: Author

13:40:12 13:40:24

4793

13:40:51 13:41:06 13:41:18

2616 290

Time

Result

13:40:33

2334

Clear History

NCBI | NLM | NIH
Department of Health & Human Services Write to the Help Desk

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OM nucleic - nucleic search, using sw model

Run on: March 22, 2005, 22:07:10; Search time 4250 Seconds

(without alignments)

9762.366 Million cell updates/sec

Title: US-10-653-681A-1

Perfect score: 1090

Sequence: 1 caaaaacagcaacagaaagc.....ataaaaaaaataataatcat 1090

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 34239544 seqs, 19032134700 residues

Total number of hits satisfying chosen parameters: 68479088

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : EST:*

1: gb_estl:*

2: gb_est2:*

3: gb_htc:*

4: gb_est3:*

5: gb_est4:*

6: gb_est5:*

7: gb_est6:*

8: gb_gss1:*

9: gb_gss2:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Resu	ult No.	Score	% Query Match	Length _e	DB	ID	Descripti	.on
		705.4	64.7	735	5	BM981698	ВМ981698	UI-CF-EN1
	2	685.2	62.9	1586	3	CR607509	CR607509	full-leng
С	3	666	61.1	746	2	BF688991	BF688991	602185236
С	4	652.2	59.8	796	5	BX337598	BX337598	BX337598
	5	647	59.4	909	4	BG169378	BG169378	602320937
С	6	586.6	53.8	666	1	AI924753	AI924753	wn58a02.x
	7	585	53.7	613	6	CB132708	CB132708	K-EST0183
С	8	573	52.6	595	5	BM983180	BM983180	UI-CF-EN1

С	9	570.8	52.4	593	5	BU677104	BU677104 UI-CF-DU1
	10	542.4	49.8	623	4	BM793014	BM793014 K-EST0073
	11	526.2	48.3	969	7	CO581979	CO581979 ILLUMIGEN
	12	522	47.9	1012	5	BQ943650	BQ943650 AGENCOURT
С	13	521.6	47.9	540	1	AA804597	AA804597 nk97e06.s
С	14	518.2	47.5	644	6	CA450136	CA450136 UI-CF-FN0
	15	517	47.4	588	2	BE785963	BE785963 601478213
	16	506.8	46.5	881	7	CO580792	· CO580792 ILLUMIGEN
	17	502.6	46.1	704	7	CO582646	CO582646 ILLUMIGEN
	18	498.8	45.8	620	6	CB118695	CB118695 K-EST0165
	19	481.4	44.2	635	7	CV334625	CV334625 IL3-UT011
С	20	475.4	43.6	496	2	AW379341	AW379341 MR0-HT024
С	21	460	42.2	920	2	BE964368	BE964368 601658069
	22	456.4	41.9	592	4	BM819663	BM819663 K-EST0087
	23	449.4	41.2	786	5	BQ221381	BQ221381 AGENCOURT
С	24	447.2	41.0	452	1	AI393702	AI393702 tg66d01.x
С	25	446.4	41.0	456	1	AI292337	AI292337 qm77c02.x
С	26	446.4	41.0	458	1	AI744504	AI744504 wg09a09.x
С	27	446.4	41.0	460	1	AI291463	AI291463 qm73h04.x
	28	446.4	41.0	582	5	BP278752	BP278752 BP278752
С	29	432	39.6	446	5	BX104876	BX104876 BX104876
С	30	426.6	39.1	445	1	AI301329	AI301329 qn27e09.x
	31	426.4	39.1	914	7	CO775128	CO775128 ILLUMIGEN
С	32	426	39.1	448	5	BM975664	BM975664 UI-CF-EN1
	33	417	38.3	445	6	CB161124	CB161124 K-EST0220
	34	402.4	36.9	581	5	BP263763	BP263763 BP263763
С	35	392.6	36.0	415	1	AA947514	AA947514 oq53h01.s
	36	388.8	35.7	453	4	BG197874	BG197874 RST17122
	37	384.8	35.3	912	5	BQ220848	BQ220848 AGENCOURT
С	38	383.4	35.2	405	6	C75075	C75075 C75075 Huma
C	39	378	34.7	388	1	AI831519	AI831519 wj49h11.x
	40	373	34.2	384	2	BE787870	BE787870 601479812
С	41	370.6	34.0	386	1	AI813308 ,	AI813308 wj33c01.x
С	42	365.2	33.5	388	2	BE711936	BE711936 QV2-HT069
	43	360	33.0	796	4	BG682196	BG682196 602629503
	44	351.2	32.2	1342	3	AK075865	AK075865 Mus muscu
	45	348	31.9	1236	3	AK019906	AK019906 Mus muscu

us-10-653-681a-1.n2p.rai

GenCore version 5.1.6 Copyright (c) 1993 - 2005 Compugen Ltd.

OM nucleic - protein search, using frame_plus_n2p model

March 19, 2005, 00:09:26; Search time 40.5 Seconds Run on:

(without alignments)

4018.147 Million cell updates/sec

US-10-653-681A-1 Title:

Perfect score: 1928

1 caaaaacagcaacagaaagc.....ataaaaaaaataataatcat 1090 Sequence:

Scoring table: BLOSUM62

Xgapop 10.0 , Xgapext 0.5 Ygapop 10.0 , Ygapext 0.5 Fgapop 6.0 , Fgapext 7.0 Delop 6.0 Delext

Searched:

513545 segs, 74649064 residues

Total number of hits satisfying chosen parameters:

1027090

Minimum DB seq length: 0 Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Command line parameters:

-MODEL=frame+_n2p.model -DEV=xlp -q=/cgn2_1/USPTO_spool_p/US10653681/runat_18032005_171051_1302/app_query.fasta_1.128

-DB=Issued_Patents_AA -QFMT=fastan -SUFFIX=n2p.rai -MINMATCH=0.1 -LOOPCL=0 -LOOPEXT=0 -UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi

-LIST=45 -DOCALIGN=200 -THR_SCORE=pct -THR_MAX=100 -THR_MIN=0 -ALIGN=15

-MODE=LOCAL -OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=2000000000 -USER=US10653681_@CGN_1_1_46_@runat_18032005_171051_1302 -NCPU=6 -ICPU=3 -NO_MMAP -LARGEQUERY -NEG_SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG -DEV_TIMEOUT=120 -WARN_TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6

-FGAPEXT=7 -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7

Database : Issued_Patents_AA:*

/cgn2_6/ptodata/1/iaa/5A_COMB.pep:*
/cgn2_6/ptodata/1/iaa/5B_COMB.pep:*
/cgn2_6/ptodata/1/iaa/6A_COMB.pep:*
/cgn2_6/ptodata/1/iaa/6B_COMB.pep:*
/cgn2_6/ptodata/1/iaa/PCTUS_COMB.pep:*
/cgn2_6/ptodata/1/iaa/backfiles1.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Result No.	Score	% Query Match	Length	DB	ID	Description
1	1265	65.6	356	4	US-09-949-016-7215	Sequence 7215, Ap

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us-10-653-681a-1.n2p.rai
                                                                                                                           Sequence 340, App
             1154
                             59.9
                                              316
                                                                US-09-702-705-340
                             59.9
                                                                US-09-736-457-340
             1154
                                              316
                                                         4 US-09-614-124B-340
4 US-09-671-325-340
4 US-09-589-184-340
4 US-09-658-824-340
                                              316
             1154
                            59.9
             1154
1154
                            59.9
59.9
                                              316
                                               316
                            59.9
                                              316
             1154
               889
                            46.1
                                              316
                                                         3 US-08-801-344-4
                                                                                                                           Sequence 4, Appli
                                                        3 US-09-498-599-4
1 US-08-585-595-3
4 US-09-270-767-45294
                                              316
316
                                                                                                                           Sequence 4, Appli
               889
                            46.1
                                                                                                                      Sequence 3, Appli
Sequence 45294, A
               877
                            45.5
10
               612
                                              302
11
                            31.7
                                                        4 US-09-949-016-6679
4 US-09-949-016-11456
4 US-09-949-016-6133
                                                                                                                           Sequence 6679, Ap
Sequence 11456, A
Sequence 6133, Ap
Sequence 7170, Ap
          586.5
585.5
532
                                              326
347
323
12
13
                            30.4
                            30.4
27.6
27.5
                                              325
323
                                                         4 US-09-949-016-7170
15
               530
                                                        4 US-09-949-010-7170
4 US-09-702-705-1821
4 US-09-671-325-1821
3 US-08-532-896-2
4 US-09-643-597-172
4 US-09-480-884A-172
                                                                                                                           Sequence 1821, Ap
                            27.3
16
               526
                                                                                                                           Sequence 1821, Ap
               526
526
                            27.3
27.3
                                              323
323
323
364
17
                                                                                                                       Sequence 1821, Ap
Sequence 1821, Ap
Sequence 2, Appli
Sequence 172, App
Sequence 173, App
18
               516
                            26.8
19
20
21
22
               514
                            26.7
26.7
                                              364
               514
                            26.7
                                              364
                                                        4 US-09-702-705-783
               514
                                              364 4 US-09-736-457-783
364 4 US-09-542-615A-172
                                                                                                                           Sequence 783, App
23
               514
                            26.7
                                                                                                                       Sequence 173, App
Sequence 172, App
24
25
               514
                            26.7
                                                                                                                          Sequence 172, App
Sequence 783, App
Sequence 783, App
Sequence 783, App
Sequence 172, App
                                                        4 US-09-606-421B-172
               514
                                              364
                            26.7
                                                        4 US-09-614-124B-783
4 US-09-671-325-783
4 US-09-589-184-783
26
27
28
29
               514
                            26.7
                                              364
                            26.7
26.7
26.7
                                              364
364
               514
               514
               514
                                               364
                                                         4 US-09-466-396A-172
                                                                                                                           Sequence 172, App
30
                            26.7
                                              364
                                                         4 US-09-476-496A-172
               514
                                                                                                                          Sequence 1/2, App
Sequence 172, App
Sequence 172, App
Sequence 9693, Ap
Sequence 2, Appli
Sequence 7078, Ap
                                                        4 US-09-470-490A-172

4 US-09-630-940B-172

4 US-09-658-824-783

4 US-09-285-479-172

4 US-09-949-016-9693

3 US-08-853-839-2

4 US-09-949-016-7078
               514
514
                                              364
364
31
                            26.7
                            26.7
32
33
                                              364
297
323
283
309
309
309
               514
                             26.7
               485
473
459
34
35
36
37
                            25.2
24.5
                            23.8
                                                                                                                          Sequence 70/8, Ap
Sequence 2, Appli
Sequence 2, Appli
Sequence 2, Appli
Sequence 46713, A
Sequence 14, Appl
Sequence 12, Appl
Sequence 10, Appl
                            22.7
22.7
22.7
                                                               US-09-166-412-2
US-09-347-803-26
US-08-731-3208-2
               438
                                                         3
38
               438
                                                         3
                                                         4
39
               438
                                                       4 US-08-731-320B-2 Sequence 2, Appli 4 US-09-166-438-2 Sequence 2, Appli 4 US-09-270-767-46713 Sequence 46713, A 3 US-09-347-803-12 Sequence 12, Appl 3 US-09-347-803-10 Sequence 10, Appl 4 US-09-270-767-43738 Sequence 43738, A
                                               309
40
                438
                             22.7
                                               346
309
           435.5
424.5
                             22.6
41
42
                             22.0
                                               308
43
           411.5
                             21.3
                             20.9
                                               308
                402
44
           400.5
                             20.8
                                               326
```

us-10-653-681a-1.n2p.rapb

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OM nucleic - protein search, using frame_plus_n2p model March 19, 2005, 00:25:42; Search time 162 Seconds Run on: (without alignments) 4448.023 Million cell updates/sec Title: US-10-653-681A-1 Perfect score: 1928 1 caaaaacagcaacagaaagc.....ataaaaaaaataataatcat 1090 Sequence: Scoring table: BLOSUM62 Xgapop 10.0 , Xgapext 0.5 Ygapop 10.0 , Ygapext 0.5 Fgapop 6.0 , Fgapext Delop 6.0 , Delext 7.0 1401741 segs. 330541175 residues Searched: 2803482 Total number of hits satisfying chosen parameters: Minimum DB seq length: 0 Maximum DB seq length: 2000000000 Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries Command line parameters: -MODEL=frame+_n2p.model -DEV=xlp -Q=/cgn2_1/USPTO_spool_p/US10653681/runat_18032005_171053_1387/app_query.fasta_1.128 -DB=Published_Applications_AA -QFMT=fastan -SUFFIX=n2p.rapb -MINMATCH=0.1 -LOOPCL=0 -LOOPEXT=0 -UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi -LIST=45 -DOCALIGN=200 -THR_SCORE=pct -THR_MAX=100 -THR_MIN=0 -ALIGN=15 -MODE=LOCAL -OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=2000000000 -USER=US10653681_@CGN_1_1_199_@runat_18032005_171053_1387 -NCPU=6 -ICPU=3 -NO_MMAP -LARGEQUERY -NEG_SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG -DEV_TIMEOUT=120 -WARN_TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6 -FGAPEXT=7 -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7 Published_Applications_AA:* Database: /cgn2_6/ptodata/2/pubpaa/US07_PUBCOMB.pep:*
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/cgn2_6/ptodata/2/pubpaa/US06_NEW_PUB.pep:*
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/cgn2_6/ptodata/2/pubpaa/US09B_PUBCOMB.pep:*
/cgn2_6/ptodata/2/pubpaa/US09C_PUBCOMB.pep:* 8: 9: 10: 11: /cgn2_6/ptodata/2/pubpaa/US09_NEW_PUB.pep:* 12: /cgn2_6/ptodata/2/pubpaa/US10A_PUBCOMB.pep:* 13: /cgn2_6/ptodata/2/pubpaa/US10B_PUBCOMB.pep:*
/cgn2_6/ptodata/2/pubpaa/US10C_PUBCOMB.pep:*
/cgn2_6/ptodata/2/pubpaa/US10D_PUBCOMB.pep:*
/cgn2_6/ptodata/2/pubpaa/US10_NEW_PUB.pep:* 14:

15: 16: 17:

```
us-10-653-681a-1.n2p.rapb

18: /cgn2_6/ptodata/2/pubpaa/US11_NEW_PUB.pep:*

19: /cgn2_6/ptodata/2/pubpaa/US60_NEW_PUB.pep:*

20: /cgn2_6/ptodata/2/pubpaa/US60_PUBCOMB.pep:*
```

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

		%			301.1.0.11.12.23	
Result		Query				
No.	Score		Length	DR	ID	Description
· 1	1268	65.8	358	9	us-09-925-299-923	Sequence 923, App
2	1268	65.8	358	10	US-09-925-299-923	Sequence 923, App
. 3	1159	60.1	316	14	US-10-274-375-3	Sequence 3, Appli
4	1154	59.9	316	9	US-09-736-457-340	Sequence 340, App
Ś	1154	59.9	316	9	US-09-902-941-340	Sequence 340, App
6	1154	59.9	316	9	US-09-849-626-340	Sequence 340, App
2 3 4 5 6 7	1154	59.9	316	10	US-09-476-300-340	Sequence 340, App
8	1154	59.9	316	14	US-10-017-754-340	Sequence 340, App
9	1154	59.9	316	14	us-10-113-872-340	Sequence 340, App
10	1154	59.9	316	15	us-10-283-017-340	Sequence 340, App
11	1073	55.7	316	14	us-10-274-375-2	Sequence 2, Appli
12	1058	54.9	316	14	us-10-274-694 - 18	Sequence 18, Appl
13	889	46.1	316	15	US-10-205-331-28	Sequence 28, Appl
14	879	45.6	316	16	US-10-451-861-30	Sequence 30, Appl
15	876	45.4	315	16	US-10-826-679-1	Sequence 1, Appli
16	875	45.4	316	16	US-10-408-765A-1573	Sequence 1573, Ap
17	875	45.4	316	16	US-10-416-330-43	Sequence 43, Appl
18	784	40.7	318	15	US-10-258-080-10	Sequence 10, Appl
19	774	40.1	245	17	US-10-653-681A-4	Sequence 4, Appli
20	638.5	33.1	176	16	US-10-451-861-31	Sequence 31, Appl
21	622	32.3	301	14	US-10-321-204-51	Sequence 51, Appl
22	613.5	31.8	302	15	US-10-038-854-44	Sequence 44, Appl
23	586.5	30.4	326	15	US-10-161-927-88	Sequence 88, Appl
24	583.5	30.3	279	15	US-10-038-854-398	Sequence 398, App
25	582.5	30.2	307	14 15	US-10-354-358-122 US-10-038-854-52	Sequence 122, App Sequence 52, Appl
26	579.5	30.1	269 325	16	US-10-036-634-32 US-10-472-317-36	Sequence 36, Appl
27 28	571 560	29.6 29.0	325	14	US-10-472-317-36 US-10-354-358-114	Sequence 114, App
28 29	560	29.0	353	9	US-09-925-301-1181	Sequence 1181, Ap
30	560	29.0	353	14	US-10-106-698-4568	Sequence 4568, Ap
31	557.5	28.9	304	14	US-10-274-694-4	Sequence 4, Appli
32	554	28.7	323	14	US-10-321-204-30	Sequence 30, Appl
33	540	28.0	323	15	US-10-038-854-14	Sequence 14, Appl
34	539	28.0	323	14	US-10-177-293-4	Sequence 4, Appli
3.5	539	28.0	323	14	US-10-321-204-1	Sequence 1, Appli
36	539	28.0	323	15	US-10-038-854-124	Sequence 124, App
\ 37	539	28.0	323	15	US-10-403-161-78	Sequence 78, Appl
38	539	28.0	323	15	US-10-403-161-94	Sequence 94, Appl
39	539	28.0	337	15	US-10-403-161-90	Sequence 90, Appl
40	537	27.9		16	US-10-476-033-2	Sequence 2, Appli
41	536	27.8	323	14	US-10-321-204-7	Sequence 7, Appli
42	536	27.8	323	15	US-10-038-854-122	Sequence 122, App
43	536	27.8	323	15	us-10-403-161-86	Sequence 86, Appl
44	536	27.8	323	15	US-10-403-161-92	Sequence 92, Appl
45	534	27.7	323	15	US-10-403-161-82	Sequence 82, Appl

OM nucleic - protein search, using frame plus n2p model

Run on: March 19, 2005, 00:08:46; Search time 49.5 Seconds

(without alignments)

4237.425 Million cell updates/sec

Title: US-10-653-681A-1

Perfect score: 1928

Sequence: 1 caaaaacagcaacagaaagc.....ataaaaaaaataataatcat 1090

Scoring table: BLOSUM62

Xgapop 10.0 , Xgapext 0.5 Ygapop 10.0 , Ygapext 0.5 Fgapop 6.0 , Fgapext 7.0 Delop 6.0 , Delext 7.0

Searched: 283416 seqs, 96216763 residues

Total number of hits satisfying chosen parameters: 566832

Minimum DB seg length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Command line parameters:

-MODEL=frame+ n2p.model -DEV=xlp

Q=/cgn2_1/USPTO_spool_p/US10653681/runat_18032005_171051_1290/app_query.fasta_1.

-DB=PIR_79 -QFMT=fastan -SUFFIX=n2p.rpr -MINMATCH=0.1 -LOOPCL=0 -LOOPEXT=0

-UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi -LIST=45

-DOCALIGN=200 -THR SCORE=pct -THR MAX=100 -THR MIN=0 -ALIGN=15 -MODE=LOCAL

-OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=2000000000

-USER=US10653681 @CGN 1 1 63 @runat 18032005 171051 1290 -NCPU=6 -ICPU=3

-NO MMAP -LARGEQUERY -NEG SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG

-DEV TIMEOUT=120 -WARN_TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6

-FGAPEXT=7 -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7

Database : PIR_79:*

1: pir1:*

2: pir2:*

3: pir3:*

4: pir4:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

		**				
Result		Query		_		
No.	Score	Match	Length D)B	ID	Description
1	1017	52.7	316	2	A53440	aldose reductase h
2	961	49.8	316	2	A37990	aldose reductase h
3	889	46.1	316	1	A60603	aldehyde reductase
4	879	45.6	316	2	149484	aldehyde reductase
5	877	45.5	315	1	A35452	aldehyde reductase
6	875	45.4	316	1	A39763	aldehyde reductase
7	870	45.1	302	2	A34406	aldehyde reductase
8	870	45.1	316	2	A59021	aldehyde reductase
9	586.5	30.4	326	1	S41120	cholestenone 5beta
10	571	29.6	325	1	JN0629	alcohol dehydrogen
11	564.5	29.3	326	1	S15835	3-oxo-5beta-steroi
12	560	29.0	325	2	A33851	alcohol dehydrogen
13	554	28.7	323	1	A45366	20alpha-hydroxyste
14	539	28.0	323	2	A53436	3-alpha-hydroxyste
15	536	27.8	323	2	JC5240	3alpha-hydroxychol
	532		323	1	A57407	chlordecone reduct
16		27.6	323	1		prostaglandin-F sy
17	527	27.3		1	JH0575	prostaglandin-F sy prostaglandin-F sy
18	524	27.2	323		A28396	chlordecone reduct
19	522	27.1	. 323	2	173676	
20	517	26.8	329	2	153872	dihydrodiol dehydr
21	516	26.8	320	2	173675	chlordecone reduct
22	514	26.7	324	1	JC4280	carbonyl reductase
23	510	26.5	323	2	173674	chlordecone reduct
24	509	26.4	323	1	A56424	estradiol 17beta-d
25	508	26.3	323	2	B57407	3alpha-hydroxyster
26	491	25.5	316	2	т26766	hypothetical prote
27	474	24.6	323	1	JC2330	luteal 20-alpha-hy
28	471.5	24.5	⁻ 322	1	A39350	3alpha-hydroxyster
29	451	23.4	309	2	A84599	hypothetical prote
30	449.5	23.3	309	2	B84599	hypothetical prote
31	433.5	22.5	327	1	S76143	probable aldehyde
32	418	21.7	290	2	T02543	aldehyde dehydroge
33	398	20.6	496	2	B89027	protein T08H10.1 [
34	396.5	20.6	284	1	A32950	probable aldehyde
35	389	20.2	310	2	T17013	D-sorbitol-6-phosp
36	386	20.0	313	2	T09670	abscisic acid acti
37	384.5	19.9	317	2	T25526	hypothetical prote
38	380	19.7	274	2	AC1535	oxydoreductases ho
39	379.5	19.7	225	1	CYFGE	epsilon-crystallin
40	377	19.6	277	2	D89964	hypothetical prote
41	377	19.6	312	1	S22846	probable aldehyde
42	374	19.4	252	2	S61515	dihydrodiol dehydr
43	373	19.3	274	2	AG1177	oxydoreductases ho
44	370	19.2	350	2	B84797	probable alcohol d
45	368.5	19.1	280	2	D69988	plant metabolite d
40	500.5	17.1	200	_	203200	Linno meanering a

us-10-653-681a-1.n2p.rup

GenCore version 5.1.6 Copyright (c) 1993 - 2005 Compugen Ltd.

OM nucleic - protein search, using frame_plus_n2p model

March 19, 2005, 00:08:07; Search time 220.5 Seconds Run on:

(without alignments)

5062.733 Million cell updates/sec

Title: US-10-653-681A-1

Perfect score: 1928

1 caaaaacaqcaacaqaaaqc.....ataaaaaaaataataatcat 1090 Seauence:

Scoring table: BLOSUM62

Xgapop 10.0 , Xgapext 0.5 Ygapop 10.0 , Ygapext 0.5 Fgapop 6.0 , Fgapext 7.0 6.0 , Delext Delop

1612378 segs, 512079187 residues Searched:

Total number of hits satisfying chosen parameters: 3224756

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Command line parameters:

-MODEL=frame+_n2p.model -DEV=xlp -Q=/cgn2_1/USPTO_spool_p/US10653681/runat_18032005_171050_1279/app_query.fasta_1.128

-DB=UniProt_03 -QFMT=fastan -SUFFIX=n2p.rup -MINMATCH=0.1 -LOOPCL=0 -LOOPEXT=0 -UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi -LIST=45 -DOCALIGN=200 -THR_SCORE=pct -THR_MAX=100 -THR_MIN=0 -ALIGN=15 -MODE=LOCAL

-OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=2000000000 -USER=US10653681_@CGN_1_1_244_@runat_18032005_171050_1279 -NCPU=6 -ICPU=3 -NO_MMAP -LARGEQUERY -NEG_SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG -DEV_TIMEOUT=120 -WARN_TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6 -FGAPEXT=7 -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7

Database: UniProt_03:*

> 1: uniprot_sprot:* uniprot_trembl:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Result No.	Score	% Query Match L	ength (DB 	ID .	Description
1	1159	60.1	316	2	AKBA_HUMAN	060218 homo sapien
2	1149	59.6	316		Q8IWZ1	Q8iwz1 homo sapien
3	1029	53.4	315		ALD2_CRIGR	008782 cricetulus
4	1013	52.5	315		ALD2_MOUSE	P45377 mus musculu
5	1001	51.9	316		Q91W30	Q91w30 rattus norv

				us	-10-653-681a-1.n2p.rup		
6	992	51.5	316	2	Q8BIV6	o8biv6	mus musculu
6 7	990	51.3	316		Q8CI22		mus musculu
8	983	51.0	316	` <u> </u>	Q6AY99		rattus norv
9	965	50.1	315		ALD1_MOUSE		mus musculu
10	961	49.8	303	1 2 2	Q68EJ1		rattus norv
11	938	48.7	316	2	Q9QZI2		rattus norv
12			315	1			oryctolagus
	898	46.6		1	ALDR_RABIT		
13	884	45.9	315	1	ALDR_RAT		rattus norv
14	877	45.5	315	1	ALDR_MOUSE		mus musculu
15	876	45.4	315	1	ALDR_HUMAN		homo sapien
16	875	45.4	316	2	Q6FGA4		homo sapien
17	873	45.3	315	1	ALDR_BOVIN	P16116	bos taurus
18	871	45.2	315	1	ALDR_PIG		sus scrofa
19	870	45.1	316	2	Q6ICP2	Q6icp2	
20	859	44.6	317	2	Q90w83	Q90w83	gallus gall
21	839	43.5	327	2	Q6AZC3	Q6azc3	brachydanio
22	812	42.1	318	2 2 2 2 2 2 2 2 1	Q6PAB5	Q6pab5	xenopus lae
23	801	41.5	315	2	Q6IQU1	06iau1	brachydanio
24	729.5	37.8	319	2	Q9DDC7		lepidodacty
25	659	34.2	316	5	Q9VTK9		drosophila
26	654.5	33.9	350	5	Q8IQF8	08iaf8	drosophila
27	623	33.9 32.3	301	ī	AKE1_MOUSE	094c+1	mus musculu
28	623	32.3	324		Q6AZW2		brachydanio
29	607.5	31.5	320	5	Q96JD6		homo sapien
30	603	31.3	327	5	Q6GMC7		xenopus lae
31	593	30.8	325	2 2 2 2	Q8VCX1		mus musculu
			323	1		051057	homo canion
32	586.5	30.4	326	1	AKD1_HUMAN	P31037	homo sapien
33	582.5	30.2	307	2	Q9BU71		homo sapien
34	582.5	30.2	326	2 2 2 1	Q9TV64	Q9TV64	oryctolagus
35	581 <u>.5</u>	30.2	335	2	Q7PCV3	Q/pcv3	anopheles g
36	577	29.9	292	2	Q7PF06	Q/p†06	anophelesg
37	576	29.9	324	1	AKA1_MOUSE		mus muscu <u>l</u> u
38	572	29.7	325	2	Q80XJ7	Q80xj7	mus musculu
39	571	29.6	324	1	AKA1_RAT		rattus norv
40	564.5	29.3	326	1	AKD1_RAT		rattus norv
41	560	29.0	324	1	AKA1_HUMAN		homo sapien
42	555	28.8	333	2	Q6DKM7		xenopus lae
43	554	28.7	323	1	PE2R_RABIT	P80508	oryctolagus
44	550.5	28.6	304	2	Q6P702		xenopus lae
45	546	28.3	324	ī	AKA1_PIG		sus scrofa
	5.5			_	· ·· · · · · · · · · · · · · · · · · ·		

OM nucleic - nucleic search, using sw model

March 22, 2005, 19:48:55; Search time 5146 Seconds Run on:

(without alignments)

10263.542 Million cell updates/sec

US-10-653-681A-1 Title:

Perfect score: 1090

1 caaaaacagcaacagaaagc.....ataaaaaaaataataatcat 1090 Sequence:

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

4708233 segs, 24227607955 residues Searched:

Total number of hits satisfying chosen parameters: 9416466

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

GenEmbl:* Database :

1: qb ba:*

2: gb htg:*

3: gb in:*

4: gb om:*

5: gb ov:*

6: gb pat:*

7: gb_ph:*
8: gb_pl:*
9: gb_pr:*
10: gb_ro:*

11: gb_sts:*

12: gb_sy:*

13: gb_un:*

14: gb_vi:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Result No.	Score	% Query Match	Length	DB	ID	 Description
1 2	833 830.8				HSU37100 CQ718316	U37100 Homo sapien CQ718316 Sequence

	3	828.2	76.0	1551	9	BC008837		BC008837	Homo sapi
	4	828.2	76.0	1560	6	CQ776685			Sequence
	5	820.2	75.2	1611	9	AF524864		AF524864	Homo sapi
	6	796.6	73.1	1316	6	AR272611		AR272611	Sequence
	7	796.6	73.1	1316	6	AR276192	·	AR276192	Sequence
•	8	796.6	73.1	1316	6	AR406467		AR406467	Sequence
	9	796.6	73.1	1316	6	AR440317		AR440317	Sequence
	10	796.6	73.1	1316	6	AR472475		AR472475	Sequence
	11	796.6	73.1	1316	6	AR543128		AR543128	Sequence
	12	796.6	73.1	1316	6	AX062696			Sequence
	13	796.6	73.1	1316	6	AX367613			Sequence
	14	796.6	73.1	1316	9	AF052577		AF052577	Homo sapi
	15	718	65.9	1315	6	AX743782		AX743782	Sequence
	16	651.8	59.8	3994	9	AL669847		AL669847	Human DNA
	17	651.8	59.8	121210	9	AL607022		AL607022	Human DNA
	18	574	52.7	574	9	AF044961			Homo sapi
	19	508	46.6	1080	6	AX772965			Sequence
	20	459.4	42.1	951	9	BT006794			Homo sapi
	21	459.4	42.1	951	12	BT007750			0 Synthetic
	22	458.4	42.1	948	9	CR541801	•		Homo sapi
	23	431	39.5	951	6	AX380448			Sequence
	24	431	39.5	951	6	AX772962			Sequence
	25	395.6	36.3	473	9	AY347931			Macaca ra
	26	377.8		144234	2	AP002425			Homo sapi
С	27	377.8		144279	2	AP001570			Homo sapi
С	28	377.8	34.7	216972	9	AC067819			Homo sapi
	29	374.8	34.4	585	6	CQ732993			Sequence
С	30	361		137557	9	AC005909			Homo sapi
	31	360.4	33.1	364	6	AX247463			Sequence
	32	358.4		163631	9	AC009276			Homo sapi
С	33	358.4		170919	9	AC078847			Homo sapi
С	34	358.4		177373	2	AP002452			Homo sapi
	35	358.4		196039	2	AC055757			Homo sapi
	36	357.2	32.8	1268	10				Cricetulus
	37	354.4	32.5	1400	10	BC037690			0 Mus muscu
	38	353.4	32.4	356	6	AX247461			Sequence
	39	353.4	32.4	1446	10				3 Rattus no
	40	344.6	31.6	1413	10	AF182168			8 Rattus no
	41	332	30.5	1315	10	BC005789			9 Mus muscu
	42	331.6	30.4	1304	10	MMU04204			Mus musculu
	43	327.2	30.0	1225	6	CQ777549			Sequence
	44	327.2	30.0	1225	10				Mouse vas d
	45	327	30.0	993	10	RNO277957		AJ27795	7 Rattus no

Job time : 5184 secs

us-10-653-681a-1.rng

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OM nucleic - nucleic search, using sw model

March 22, 2005, 19:42:42; Search time 701 Seconds Run on:

(without alignments)
9204.733 Million cell updates/sec

Title:

US-10-653-681A-1

Perfect score:

1090

Sequence:

1 caaaaacagcaacagaaagc.....ataaaaaaaataataatcat 1090

Scoring table: IDENTITY_NUC

Gapop 10.0 , Gapext 1.0

Searched:

4390206 seqs, 2959870667 residues

Total number of hits satisfying chosen parameters:

8780412

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database :

N_Geneseq_16Dec04:* geneseqn1980s:* geneseqn1990s:* geneseqn2000s:* geneseqn2001as:* geneseqn2001bs:* geneseqn2002as:* geneseqn2002bs:* geneseqn2003as:* geneseqn2003bs:* geneseqn2003cs:* 10: geneseqn2003ds:* 11: genesegn2004as:* 12: 13: geneseqn2004bs:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Result No.	Score	% Query Match	Length	DB	ID	Description
1	833	76.4	1337	5	AAS68608	Aas68608 DNA encod
2	833	76.4	1337	10	ADD71032	Add71032 Human ald
3	832.2	76.3	1508	3	AAC98140	Aac98140 Human col
4	828.2	76.0	1560	12	ADJ75119	Adj75119 Marker ge
5	828.2	76.0	1560	12	ADN04246	Adn04246 Antipsori
6	828.2	76.0	1560	13	ACN38728	Acn38728 Tumour-as
7	828.2	76.0	1560	13	ADS85007	Ads85007 Human ato
8	822.8	75.5	1549	12	ADK70274	Adk70274 Respirato
ğ	796.6	73.1	1316	5	AAF68405	Aaf68405 Human lun
10	796.6	73.1	1316		ABK38316	Abk38316 cDNA enco

```
us-10-653-681a-1.rng
                                                                                                            Ads73134 Human kid
Aca10645 Human lun
Abx99596 Lung canc
Adh45842 Human lun
              796.6
                                                          ADS73134
                             73.1
                                          1316
                             73.1
73.1
73.1
73.1
     12
13
                                                     8
              796.6
                                          1316
                                                           ACA10645
              796.6
                                           1316
                                                           ABX99596
              796.6
                                                      10 ADH45842
     14
                                           1316
              796.6
                                                             ADE72379
                                                                                                              Ade72379 Human lun
     15
                                           1316
                                                      12
              796.6
                             73.1
70.7
     16
                                           1316
                                                      13
                                                             ADJ19761
                                                                                                              Adj19761 Human lun
                                                                                                              Adh13722 Human ENZ
                                                             ADH13722
ADC97771
              770.8
                                           1621
     17
                                                      12
                                                                                                             Adc97771 Human ARL
Acn92921 Breast ca
Adr98739 Lung spec
Adh45334 Human enz
                             65.9
                                           1315
                                                      10
     18
                718
                                          1816
770
1170
              635.4
                             58.3
56.5
                                                             ACN92921
ADR98739
                                                      11
     19
                                                      13
12
     20
                  616
                                                                                                           Adh45334 Human enz
Acc83986 Human ald
Abz84625 Toxicolog
Adc10183 Human NOV
Aba94733 Human NOV
Aba94733 Human NOV
Aas39335 Novel hum
Aas39333 Novel hum
Aas72230 DNA encod
Aas92672 DNA encod
Adj75983 Marker ge
Adp28822 Human sec
Adj75983 Marker ge
Adp28822 Human lun
Adc5831 Human lun
Abq92236 Human lun
Add28651 Human lun
Add36746 Human lun
Adm56549 Human lun
Adm86549 Human lun
Adm89593 Human lun
Ada28650 Human lun
Ada28650 Human lun
Ada28650 Human lun
                                                           ADH45334
              540.2
                             49.6
     21
              508
439.8
                                                     9 ACC83986
10 ABZ84625
                             46.6
                                           1080
     22
                                            558
971
                             40.3
     23
     24
                  432
                             39.6
                                                      10 ADC10183
                  431
                                            951
                                                      6 ABA94733
     25
                             39.5
     26
                  431
                                                     10 ADC10185
                             39.5
                                            966
                                                     4 AAS39335
4 AAS39333
5 AAS72230
5 AAS92672
     27
28
29
30
              360.4
                             33.1
                                            364
              353.4
333.8
                                          356
1926
                             32.4
                             30.6
              333.8
                                           1926
                             30.6
                                           3620
1225
540
585
585
585
              333.8
     31
                             30.6
                                                      5 AAS69995
                                                           ADJ75983
     32
              327.2
317
                                                     12
                             30.0
                             29.1
                                                      12 ADP28822
     33
                             29.0
29.0
29.0
     34
                  316
                                                      2
                                                            AAZ24592
                                                      3 AAC65831
6 ABL49050
     35
                  316
     36
37
                   316
                                             585
585
                                                      6 ABQ92236
9 ADA28651
                             29.0
                  316
                316
                             29.0
     38
                                             585
585
                                                      10 ADE53611
     39
                  316
                             29.0
                             29.0
                                                      10
                                                            ADH36746
     40
                  316
                                            585
585
857
858
858
                                                             ADM56549
     41
                  316
                              29.0
                                                      12
                                                            ADN89593
     42
                  316
                             29.0
                                                      12
                                                                                                             Ada28650 Human lun
     43
                  316
                             29.0
                                                      9
                                                            ADA28650
C
                              29.0
                                                      2
                                                            AAZ24591
                                                                                                            Aaz24591 Human lun
     44
                   316
                  316
                             29.0
                                                     3 AAC65830
                                                                                                             Aac65830 Human lun
```

us-10-653-681a-1.rni

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OM nucleic - nucleic search, using sw model

March 22, 2005, 22:13:45; Search time 228 Seconds Run on:

(without alignments) 7822.551 Million cell updates/sec

Title: US-10-653-681A-1

Perfect score: 1090

1 caaaaacagcaacagaaagc.....ataaaaaaaataataatcat 1090 Sequence:

Scoring table: IDENTITY_NUC

Gapop 10.0 , Gapext 1.0

1202784 seqs, 818138359 residues Searched:

Total number of hits satisfying chosen parameters: 2405568

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database: Issued_Patents_NA:*

/cgn2_6/ptodata/1/ina/5A_COMB.seq:*
/cgn2_6/ptodata/1/ina/5B_COMB.seq:*
/cgn2_6/ptodata/1/ina/6A_COMB.seq:*
/cgn2_6/ptodata/1/ina/6B_COMB.seq:*
/cgn2_6/ptodata/1/ina/PCTUS_COMB.seq:*
/cgn2_6/ptodata/1/ina/backfiles1.seq:* 2: 3: 4:

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

		%	•			
Result		Query				Baranintian
No.	Score	Match	Length		ID	Description
1	828.2	76.0	1515	4	us-09-949-016-1344	Sequence 1344, Ap
2	796.6	73.1	. 1316	4	us-09-702 - 705-323	Sequence 323, App
3	796.6	73.1	1316	4.	us-09-736-457-323	Sequence 323, App
4	796.6	73.1	1316	4	US-09-614-124B-323	Sequence 323, App
5	796.6	73.1	1316	4	us-09-671-325-323	Sequence 323, App
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7	796.6	73.1	1316	4	us-09-658-824-323	Sequence 323, App
8	389.2	35.7	914	4	us-09-949-016-3127	Sequence 3127, Ap
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10	328.6	30.1	601	4	us-09-949-016-46452	Sequence 46452, A
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14	316	29.0	585	4	us-09-542-615A-92	Sequence 92, Appl
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us-10-653-681a-1.rni
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Sequence 3, Appli
Sequence 3, Appli
Sequence 46451, A
Sequence 13724, A
Sequence 14649, A
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4 US-09-270-767-13724
4 US-09-949-016-14869
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             288.4
170.2
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                                       601
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Sequence 31, Appl
Sequence 6562, Ap
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us-10-653-681a-1.rnpb

GenCore version 5.1.6 Copyright (c) 1993 - 2005 Compugen Ltd.

OM nucleic - nucleic search, using sw model

March 23, 2005, 00:12:25; Search time 748 Seconds Run on:

(without alignments) 8675.151 Million cell updates/sec

Title: US-10-653-681A-1

Perfect score: 1090

1 caaaaacagcaacagaaagc.....ataaaaaaaataataatcat 1090 Sequence:

Scoring table: IDENTITY_NUC

Gapop 10.0 , Gapext 1.0

5544816 segs, 2976611598 residues Searched:

Total number of hits satisfying chosen parameters: 11089632

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0% Maximum Match 100%

Listing first 45 summaries

Published_Applications_NA:* Database :

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/cgn2_6/ptodata/2/pubpna/US06_NEW_PUB.seq:*
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/cgn2_6/ptodata/2/pubpna/US09_NEW_PUB.seq:* 8: 9: 10: 11: 12: /cgn2_6/ptodata/2/pubpna/US10A_PUBCOMB.seq:* 13: /cgn2_6/ptodata/2/pubpna/US10B_PUBCOMB.seq:*/cgn2_6/ptodata/2/pubpna/US10C_PUBCOMB.seq:* 14: 15: 16: 17:

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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Result No.	Score	% Query Match	Length	DB	ID	Description
1	1090	100.0	1090	19	US-10-653-681A-1	Sequence 1, Appli

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us-10-653-681a-1.rnpb
                                                                                                                                                                               Sequence 150, App
Sequence 150, App
Sequence 323, App
                                                                                                                                              1508
                                                 832.2
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                                                 796.6
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13
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707
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353.4
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                                                               292
                                                               292
                                                                                                     26.8
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```

OM nucleic - protein search, using frame plus n2p model March 18, 2005, 23:59:48; Search time 182.5 Seconds Run on: (without alignments) 4619.932 Million cell updates/sec US-10-653-681A-1 Title: Perfect score: 1928 1 caaaaacagcaacagaaagc.....ataaaaaaaataataatcat 1090 Sequence: Scoring table: BLOSUM62 Xgapop 10.0 , Xgapext 0.5 Ygapop 10.0 , Ygapext 0.5 Fgapop 6.0, Fgapext 7.0 6.0 , Delext Delop 2105692 segs, 386760381 residues Searched: 4211384 Total number of hits satisfying chosen parameters: Minimum DB seg length: 0 Maximum DB seq length: 2000000000 Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries Command line parameters: -MODEL=frame+ n2p.model -DEV=xlp Q=/cgn2 1/USPTO_spool_p/US10653681/runat_18032005_171050_1272/app_query.fasta_1. 1287 -DB=A Geneseq 16Dec04 -QFMT=fastan -SUFFIX=n2p.rag -MINMATCH=0.1 -LOOPCL=0 -LOOPEXT=0 -UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi -LIST=45 -DOCALIGN=200 -THR SCORE=pct -THR MAX=100 -THR MIN=0 -ALIGN=15 -MODE=LOCAL -OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=2000000000 -USER=US10653681 @CGN 1 1 224 @runat 18032005 171050 1272 -NCPU=6 -ICPU=3 -NO MMAP -LARGEQUERY -NEG SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG -DEV TIMEOUT=120 -WARN_TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6 -FGAPEXT=7 -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7 Database : A Geneseq 16Dec04:* 1: geneseqp1980s:* 2: geneseqp1990s:* 3: geneseqp2000s:* geneseqp2001s:* 4:

Pred. No. is the number of results predicted by chance to have a

5: geneseqp2002s:*

geneseqp2003as:*

geneseqp2003bs:*
geneseqp2004s:*

6:

7:

score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

		8				
Result		Query				
No.	Score	Match	Length	DB	ID	Description
						21 -04401 Wass 1 have
1	1274	66.1	339	4	ABG04421	Abg04421 Novel hum
2	1268	65.8	358	3	AAB53383	Aab53383 Human col
3	1159	60.1	316	3	AAB10872	Aab10872 Human ARL
4	1159	60.1	316	7	ABR62325	Abr62325 Human ald
5	1159	60.1	316	7	ADC97773	Adc97773 Human ARL
6	1159	60.1	316	8	ADK70537	Adk70537 Respirato
7	1159	60.1	316	8	ADJ75577	Adj75577 Marker ge
8	1159	60.1	316	8	ADN04247	Adn04247 Antipsori
9	1159	60.1	316	8	ABM80941	Abm80941 Tumour-as
10	1159	60.1	316	8	ADS85008	Ads85008 Human ato
11	1154	59.9	316	4	AAB76864	Aab76864 Human lun
12	1154	59.9	316	5	AAU85519	Aau85519 Clone #18
13	1154	59.9	316	6	ABU69491	Abu69491 Human lun
14	1154	59.9	316	6	ABU66393	Abu66393 Lung canc
15	1154	59.9	316	7	ADH45859	Adh45859 Human lun
16	1154	59.9	316	8	ADE71611	Ade71611 Human lun
17	1154	59.9	316	8	ADJ19778	Adj19778 Human lun
18	1084.5	56.2	293	8	ADH45296	Adh45296 Human enz
19	1073	55.7	316	7	ADC97772	Adc97772 Human ARL
20	1058	54.9	316	5	ABB07530	Abb07530 Human dru
21	1058	54.9	316	7	ABR62324	Abr62324 Human ald
22	1058	54.9	316	7	ADC10186	Adc10186 Human NOV
23	1056	54.8	316	7	ADC10184	Adc10184 Human NOV
24	961	49.8	316	3	AAB10871	Aab10871 Murine MV
25	961	49.8	316	8	ADJ76302	Adj76302 Marker ge
26	920	47.7	287	8	ADH13653	Adh13653 Human ENZ
27	889	46.1	316	2	AAW69357	Aaw69357 Rat lens
28	889	46.1	316	6	ABM04798	Abm04798 Rat aldos
29	884	45.9	315	7	ADE57875	Ade57875 Rat Prote
30	884	45.9	315	7	ADE57879	Ade57879 Rat Prote
31	884	45.9	315	7	ADE57883	Ade57883 Rat Prote
32	884	45.9	315	7	ADE57887	Ade57887 Rat Prote
33	884	45.9	315	7	ADE57867	Ade57867 Rat Prote
34	884	45.9	315	7	ADE57871	Ade57871 Rat Prote
35	882	45.7	316	8	ADS85097	Ads85097 Mouse ato
36	882	45.7	328	4	ABG11687	Abg11687 Novel hum
37	879	45.6	316	5	ABB83356	Abb83356 Murine TS
38	878	45.5	316	2	AAR15425	Aar15425 Human ald
39	876	45.4	315	.7	ADE57869	Ade57869 Human Pro
40	876	45.4	315	7	ADE57877	Ade57877 Human Pro
41	876	45.4	315	7	ADE57885	Ade57885 Human Pro
42	876	45.4	315	7	ADE57881	Ade57881 Human Pro
43	876	45.4	315	7	ADE57889	Ade57889 Human Pro
44	876	45.4	315	7	ADE57873	Ade57873 Human Pro
45	876	45.4	315	8	ADP90915	Adp90915 Human ald